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Solvent compositions for use in pressure-sensitive capying paper

 A solvent composition for use in pressure-sensitive copying papers comprises a mixture of vegetable oil and a mono- or di-functional ester of a fatty acid or other acid composed of a non-aromatic saturated or unsaturated straight or branched hydrocarbon chain and a single terminal carboxyl group.

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as carbonless copying paper. This invention relates to solvent compositions for use in pressure-sensitive copying paper, also known

known as "CFB" sheets) are provided, each of which is coated on its lower surface with microcapsules and ransfer type. A business forms set using the transfer type of pressure-sensitive copying paper comprises an upper sheet (usually known as a "CB" sheet) coated on its lower surface with microcapsules containing on its upper surface with colour developer composition. Imaging pressure exerted on the sheets by writing a colour former) and a lower sheet (usually known as a "CF" sheet) coated on its upper surface with a a solution in an oil solvent or solvent composition of at least one chromogenic material (alternatively termed sets. Various types of pressure-sensitive copying paper are known, of which the most widely used is the releasing or transferring chromogenic material solution on to the colour developer composition and giving typing or impact printing (e.g. dot matrix or daisy-wheel printing) ruptures the microcapsules, thereby colour developer composition. If more than one copy is required, one or more intermediate sheets (usually Pressure-sensitive copying paper is well-known and is widely used in the production of business forms

as dispersed droplets in a continuous pressure-rupturable matrix instead of being contained within discrete In a variant of the above-described arrangement, the solution of chromogenic material may be present rise to a chemical reaction which develops the colour of the chromogenic material and so produces a copy

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8 system, microcapsulos and colour developing co-reactant material are coated onto the same surface of a pressure-rupturable microcapsules. material on the sheet to produce a coloured image. rupture and release the solution of chromogenic material, which then reacts with the colour developing sheet, and writing or typing on a sheet placed above the thus-coated sheet causes the microcapsules to In another type of pressure-sensitive copying system, usually known as a self-contained or autogeneous

33 described above have typically been products of the petrochemical industry for example partially hydroge nated terphenyls, alkyl naphthalenes, diarylmethane derivatives, dibenzyl benzene derivatives or chlorinated paraffins. These "prime solvents" are usually mixed with cheaper diluents or extenders such as kerosene, The solvents used to dissolve the chromogenic materials in pressure-sensitive copying papers

ខ principle an alternative to the use of petrochemical-based solvent compositions. However, to the best of our papers, even though proposals for use of vegetable oil solvents go back many years, see for example U.S. knowledge, there has been no commercial utilization of vegetable oil solvents in pressure-sensitive copying which although of lesser solvating power, give rise to more cost-effective solvent compositions. Vegetable oils have been disclosed as solvents for use in pressure-sensitive copying papers, and are in

35 compositions for pressure-sensitive copying paper which comprise a blend of an aromatic hydrocarbon with composition which actually contains a vegetable oil. castor oil, soybean oil and corn oil, but there is no exemplification or explicit disclosure of any solvent Patents No. 2712507; 2730457 and 3016306 additionally contain an "inert diluent". The examples given of such a diluent include vegetable oils such as specified aliphatic acid diesters. European Patent Application No. 24888A discloses also that the blend may European Patent Application No. 24898A and British Patent No. 1526353 each disclose solvent

for example US Patent No. 3016308 referred to above. solvents or pigment-suspending media for pressure-sensitive copying paper has also been proposed, see The use of phthalates, for example dibutyl phthalate, and certain other esters, for example maleates, as

it is specifically directed to the use of vegetable, animal or mineral oil solvents in pressure-sensitive copying and, especially, in European Patent Application No. 262569A. The last-mentioned is of particular interest as be found, for example, in European Patent Applications Nos. 86636A (page 4), 155593A (page 11), 2343944 paper. In contrast, the references to vegetable oil solvents in the other patents just referred to were generally made in passing, the main subject of the patent not being concerned with solvent compositions at More recent disclosures of the use of vegetable oil solvents in pressure-sensitive copying paper are to

or carbinot derivative type would be a significant drawback to the use of vegetable oil solvents requirement for the replacement of tried and tested phthalide and fluoran chromogenic materials chromogenic materials which have hitherto been the most widely used chromogenic materials in the art. A dyes are preferably carbinols or C1 to C4 alkoxy derivatives of carbinols. Such carbinols or carbinol elatively unproven, or at least less well-established, chromogenic materials of the triphenylmethane carbinol derivatives differ from the phthalide chromogenic materials, e.g. Crystal Violet Lactone ("CVL") and fluoran materials in conjunction with the vegetable, animal or mineral oils disclosed. These triphenylmethane leucc European Patent Application No. 262569A requires the use of triphenylmethane leuco dye chromogenic

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An important consideration in our evaluation of vegetable oil solvents has therefore been that these

present no serious problems when used with vegetable oil solvents, either as regards solubility or colour and fluoran type. We have found that most of the widely-used phthalide and fluoran chiernogenic materials generating capability. However we did carcounter one or more of the following problems: solvents should be capable of satisfactory use with well-established chromogenic materials of the phthalide

 Wido Primary Droplet Size Distribution on Emulsification droplets in this emulsion is a key parameter in determining the size of the final microcapsules. Wide In order to encapsulate the oils, they must first be emulsified in an aqueous medium. The size of the

coating machine, the capsule coated (CB) surface of each ply within the reel is in close contact with the coloured spots and in general discolouration in CFB paper, since in a wound reel of CFB from the the case of excessively large microcapsules. These are particularly prone to damage and accidental variations in primary droplet size, and hence in microcapsule size, are disadvantageous, particularly in sensitive copying sets, where CB and CF surfaces are also in contact. colour developer (CF) surface of the adjacent ply. Spot fermation can also occur in finished pressureretained by the microcapsule walls and therefore can escape prematurely). This results in production of rupture, and may also be more permeable than smaller capsules (i.e. the capsule contents are less well

chromogenic material solution in a spherical droplet is proportional to the cube of the radius of the droplet, and that what may seem to be a relatively minor oversizing can have very significant effects in In considering the problems just described, it should be borne in mind that the volume of

colouration (see below). A wide primary droplet size distribution can also exacerbate the problem of post-printing dis-

2. Post-Printing Discolouration

forms sets, a certain amount of microcapsule damage tends to occur; and this results in release of chromogenic material solution which can transfer to an adjacent CF surface and produce discolouration Discolouration on Storage (or "post-print blacking", or "post-print blueing", depending on the colour of the copy image). as a result of formation of many small coloured specks. This is known as "post-printing discolouration" When CB and CFB papers are subjected to a printing process as part of the production of business

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reasons for this include the presence in the microcapsule coating of a small proportion of unenin the case of a blue-copy product) and is referred to generally as discolouration on storage. colour developer coating on the top surface. The effect is primarily seen as an overall greying (or blueing free chromogenic material solution can potentially migrate up through the paper and into contact with the the microcapsule walls, and premature capsule damage as a result of the strains imposed by reel tensions, or by the weight of higher cheets in the case of stacked sheeted products. In each case, the capsulated chromogenic material solution, gradual permeation of chromogenic material solution through It is found that CFB paper sometimes tends to discolour gradually on storage prior to use. The

a mono- or di-functional ester of certain organic acids. also that an improved copy intensity can be obtained, if the vegetable oil solvent is used in conjunction with It has now been found that the above-described problems can be eliminated or at least reduced, and ដូ

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paper and comprising a vegotable oil, characterized in that the solvent composition also comprises a addition to the carboxyl carbon atom). The carboxyl group is preferably a terminal carboxyl group. proportion of a mono-or di-functional ester of a non-aromatic mone-carboxylic acid having a saturated or unsaturated straight or branched hydrocarbon chain with at least three carbon atoms in the chain (i.e. in Accordingly, the present invention provides a solvent composition for use in pressure-sensitive copying

pressure-rupturable barrier. just defined, either contained in microcapsules or otherwise present in the form of isotated droplets in a The invention also extends to pressure-sensitive copying paper comprising a solvent composition 8

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at room tomporatures, but this does not matter provided that they are used with an ester with which the oil oil, soya bean oil, sunflower oil or corn oil is preferred. Certain of the oils just listed are solid or semi-solid cottonseed oil, safflowor oil, linseed oil, castor oil, babassu oil, tung oil, jojoba oil or oilicica oil. Rapeseed sunllower ell, soybean ell, cern ell, cecenut ell, palm kernel ell, palm ell, elive ell, greundnut ell, sesame ell will form a liquid blend of a workable viscosity. The vegotable oil may be any of the commonly-available vegetable oils, for example rapeseed oil,

S available, see for example "Kirk-Othmer Encyclopedia of Chemical Technology", third Edition, Volume 23 (section on "Vogetable Oits") and Volume heta (section on "Fats and Fatty Oits"), published by John Wiley hetainformation on the chemical composition, extraction, refining and purification of vegetable oils is widely

The ester used in the present solvent composition is preferably an ester of a fatty acid, i.e. an ester of

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Edition, rovised by N. Irving Sax and Richard J. Lewis, Sr. published by Van Nostrand Reinhold Company single terminal carboxyl group, the total number of carbon atoms present (including the carboxyl group) reference books, the usage in this specification, i.e. as meaning an acid derivable from an animal or Fatty acids are composed of a saturated or unsaturated straight or branched hydrocarbon chain with vegetable oil, is consistent with the definition in "Hawley's Condensed Chemical Dictionary", an acid derivable from an animal or vegetable oil, and will horeatter be referred to for convenience as a *taty ਜਰਮ ester". Whilst the expression "fatty acid" is not always defined consistently in technical

acid such as myristic acid, capric acid, caprylic acid, stearic acid, isostearic acid, palmitic acid, or lauric of coconut acid have chain lengths of 6 to 18 carbon atoms and are chiefly lauric, capric, myristic, palmitic coconut acid, i.e. a mixture of fatty acids derived from hydrolysis of coconut oil. The constituent fatty acids generally being an even number from 4 to 22. "coconutate" is also in use (it should be noted that the expression "cocoate" has no connection with and cleic acids. An ester of coconut acid will hereafter be referred to as a "cocoate", although the term acid, or of an unsaturated fatty acid such as oleic acid, or of an acid of mixed composition, for example By way of example, the fatty acid ester may be of a saturated straight or branched-chain aliphatic fath 끍

acids present in cocoa oil or cocoa butter). The ester moiety of the fatty acid or other ester used in the present solvent composition may vary

functional ester moiety is propylene glycyl (i.e. an ester moiety derived from propylene glycol) widely. For example, it may have only one carbon atom, i.e. methyl, or several carbon atoms, for example isopropy), octyl or 2-ethylhexyl. Such ester moieties are all mono-functional. An example of a suitable di-

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vegetable cits. thus a mixture of a vegetable oil and a glyceryl ester probably behaves in a manner similar to a blend of same benefits, perhaps because such esters are chemically similar to naturally-occurring tri-glycerides We have so far found that the use of a tri-functional ester such as a glyceryl ester does not give the

available products, being used in industry for a variety of applications, particularly cosmetics and other personal care products. They can be manufactured by esterification, with suitable alcohols, of fatty acids widely available. derived by relining and/or distillation of crude vegetable oils. The alcohols required for esterification are Numerous examples of mono- or di-functional esters of fatty acids as disclosed above are commercially

following, which may be used singly or in combination: Specific examples of suitable fatty acid esters for use in the present solvent composition include the

2-ethylhexyl cocoate(EHC)

methyl oleate (MO) (see note 1)

propylene glycol dicaprylate/caprate) (PGCC) (see note 2) methył isostearate (MIS)

- ć 1. "Methyl oleate" (MO) is a commercial name for a mixtue of fatty acid methyl esters in which the major monoesters having from 4 to 18 acid moiety carbon atoms (c. 10% in total) linoleate (c. 9%), methyl palmitoleate (c. 5%), methyl linolenate (c.2%) and various saturated methyl component (c. 73%) is methyl oleate but which also contains other unsaturated materials, namely methyl
- but also contains minor proportions of other acid moieties, principally lauric acid (c. 5%). 2. PGCC has caprylic acid and capric acid as the main acid moieties (c. 59% and c. 36% respectively)
- Gouda, The Netherlands. All of the above-listed esters are commercially-available, for example from Unichema International of

Of the above-listed esters, EHC and IPM are preferred.

8 used in the present solvent composition. An ester made from acid manufactured in this way is termed a natural oil but which was actually manufactured other than from a natural oil source could in principle be have actually been derived from a natural oil. However, a fatty acid which is of a kind derivable from a "synthesized fatty acid ester" In general, the acid moiety of fatty acid ester(s) suitable for use in the present solvent composition will

55 other organisms. Although they occur naturally, their commercially-available forms are generally syn naturally in secretions of certain birds and animal skins (for example in human skin), and in yeast, fungi and esters, are generally alkyl-branched esters of aliphatic carboxylic acids and aliphatic alcohols. They occur the kind found in naturally-occurring lipids may be employed. Such esters, which are often termed wax As an alternative to the use of a fatty acid ester or synthesized fatty acid ester, closely related esters of

by P.E. Kollattukudy, published by Elsevier, Amsterdam, in 1976 occurring wax ostors can be found, for example, in "Chemistry and Biochemistry of Natural Waxes", edited zolvent compositions, and is also available from Unichema International. Further information on naturally-(EHEH) is an example of a commercially-available synthesised wax ester which is usable in the present thesized from non-naturally derived alcohol and acid starting materials. 2-ethylhexyl-2-ethylhexancate

present solvent compositions, in practice curtain of them have proporties or side offects which may make them unsuitable. For example, the esters must have a workable viscosity when in a blend with the vegetable oil. Also, certain esters have an unacceptable odour (although this may have been due to impurities in the Although in principle all mono- or distunctional esters of the kind defined herein are usable in the

sample we evaluated, and would not necessarily be present in all samples). Additionally, we have found that does of course require only very simple tests or procedures, and needs no further description. Problems care must be taken to screen prospective esters for drawbacks such as just discussed. Such screening caused by the presence of undestrable impurities can of course be solved by improved purification known as a desensitizer for pressure-sensitive copying paper. Thus when seeking to work the invention developer. Again, this may well be due to the presence of impurities such as polyethylene glycol, which is and prevent or reduce proper colour development of chromogenic material on contact with colour samples of certain fatty acid esters, for example polyethyleneglycol cocceate, have a desensitizing effect

ä 8 tochnical benotits achievable by the use of the defined ester(s) have to be balanced against their high cost accommodated to a considerable extent. A further factor is that the defined esters generally have relatively poor solvating power for chromogenic materials as currently used in pressure-sensitive copying papers. compared with petrochemical-based solvents and so the relatively high cost of the defined esters can be compared with the cost of vegetable oils. However, vegetable oil solvents are generally very cheap This could potentially limit the amount of estor which can be used The relative proportions of vegetable oil and ester in the solvent composition can vary widely, but the

techniques.

the range 1:3 to 3:1 to be suitable, but these values are not to be taken as in any way indicating limits of Taking these various factors into account, we have so far found a weight ratio of vegetable oil:ester in

The present solvent composition is preferably composed substantially entirely of vegetable oil(s) and

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result of oxidation. present, for example antioxidants to counteract the well known tendency of vegetable oils to deteriorate as a In addition to the chromogenic materials dissolved in the solvent composition, other additives may be

မ္မ In use, the present solvent composition, containing dissolved chromogenic materials, is microen-

capsulated and used in conventional manner. as doscribed in U.S. Patents Nos. 2800457; 2800458; or 3041289; or by in situ polymerisation of polymer procursor material, e.g. as described in U.S. Patents Nos. 4001140; 4100103; 4105823 and 4396670. The microcapsules may be produced by coacervation of gelatin and one or more other polymers, e.g.

as 3,3-bis(4-dimethylaminophenyl)-6-dimethylaminophthalide (CVL) and 3,3-bis(1-octyl-2-methylindol-3-yl)phenylaminotluoran, or 3'-chloro-6'-cyclohexylaminotluoran; or spirobipyran derivatives such as 3'-i-propyl othyf-N-phonylamino-4'-mothyffluoran). phthalido; Ituoran dorivatives, such as 2'anilino-6'-diethylamino-3'-methyltluoran, 6'-dimethylamino-2'-(Ndibenzylamino-2,2'-spirobi-(2H-1-benzopyran). Triphenylmethyl chromogenic materials as disclosed in The chromogenic materials used in the microcapsules may be, for example, phthalide derivatives, such 2'-N-methyl-N-phenylaminofluoran-6'-N-ethyl-N(4-methyl-

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rosulting coating composition is then applied by conventional coating techniques, for example metering roll material may be, for example, wheatstarch particles or ground cellulose fibre floc or a mixture of these. The agent (or "stilt material") for protecting the microcapsules against premature microcapsule rupture. The stilt with a sultable binder, for example starch or a starch/carboxymethylcellulose mixture, and a particulate The chromogen-containing microcapsules, once produced, are formulated into a coating composition

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European Patent Application No. 262569A may also be used.

Apart from the solvent composition, the present pressure-sensitive copying paper may be conventional. Such paper is very widely disclosed in the patent and other literature, and so requires only brief further coating or air knife coating.

conventional for this type of paper, for example the thickness may be about 60 to 90 microns and the grammage about 35 to 50 g m⁻², or higher, say up to about 100 g m⁻², or even more. This grammage depends to some extent on whether the linal paper is for CB or CFB use. The higher grammages just The thickness and grammage of the present paper (before microcapsule coating) may be as is

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quoted are normally applicable only to speciality CB papers.

428994A, or German Offenlegungsshrift No. 4110354A. a phenolic resin, e.g. as described in U.S. Patent No. 3672935 or No. 4612254; or an organic acid or metal salt thereof, e.g. as described in U.S. Patent No. 3024927, European Patent Applications Nos. 2/75107A or The colour developer material used may be an acid clay, e.g. as described in U.S. Patent No. 3753761;

The invention will now be illustrated by the following Examples in which all parts, percentages and

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This illustrates the use of a solvent composition comprising rapeseed oil (RSO) and 2-ethylhexylcocoate (EHC) in 3:1 and 1:1 ratio, with a 100% rapeseed oil solvent composition as a control for comparison

ñ ericapsulation. These chromogenic materials are all commercially available and have a long history of use in in the case of the RSO/EHC compositions and 6.4% in the case of the 100% RSO composition. such as to give a black print, as is conventional in the art. The total colour former concentrations were 5.0% spirobipyran chromogen and a red bis-indolyl phthalide chromogen, and were used in relative proportions the art. They were principally CVL, a green fluoran and an orange fluoran, with smaller amounts of a blue Chromogenic materials were first dissolved in the solvent compositions to produce solutions for

25 8 measured by means of a Coulter Counter). The milling times required to achieve this median primary saving in milling minutes for the 100% RSO composition. Thus the inclusion of a proportion of EHC produces a significant solution, and the resulting dispersion was then milled to a target median droplet size of 3.2 \pm 0.2 μ m (as step of the encapsulation process, the chromogenic material solution was dispersed with stirring in gelatine carboxymethyl cellulose and vinylmethylether/maleic anhydride copolymer as anionic colloids. As an initial generally conventional gelatin coacervation technique as disclosed in British Patent No. 870476, using droplet size were 45 and 49 minutes for the 3:1 and 1:1 RSO:EHC compositions respectively, and The resulting chromogenic material solutions were encapsulated on a pilot plant scale by means of a time.

ઇ as to permit a droplet size distribution to be derived. This showed that the percentage of "oversize" inclusion of a proportion of EHC resulted in significant benefits. 1.8% for the 1:1 RSO/EHC composition and 3.5% for the 100% RSO composition. Again therefore, the droplets, defined as droplets of a size greater than 6.35 um, was 2.9% for the 3:1 RSO:EHC composition The Coulter Counter was also used to measure the percentage of droplets in different size ranges, so

spread of droplet size distribution and is the difference between the upper and lower quartile droplet sizes. The smaller the IQD value the narrower (i.e. better) the droplet size distribution. The IQD values were 1.89 ит for the 3:1 RSO:EHC composition, 1.73 ит for the 1:1 RSO:EHC composition, and 1.99 ит for the 100% RSO composition. This was corroborated by IQD calculations (IQD = Inter-Quartite Distance). IQD is a measure of the

sion was diluted with additional water and vinylmethyl ether/maleic anhydride copolymer solution to adjust the pH to about 4.2 and thereby bring about coacervation. The coacervate deposited about the emulsified oil droplets so as to form liquid-walled microcapsules. The mixture was then chilled to about added. After heating to 50-55 °C, carboxymethylcellulose solution was added. Acetic acid was then added 10°C to solidify the initially-liquid coacervate walls, after which a hardening agent The microencapsulation process was then completed in conventional manner. Specifically, the disper (glutaraidehyde) was

added to cross-link the walls and prevent their re-dissolving when the temperature rises when the chilling The resulting microcapsule dispersion was then adjusted to pH 7 with sodium hydroxide solution. operation is concluded. A further addition of vinylmethylether/maleic anhydride copolymer was then made

available 46 g m $^{-2}$ CF paper by means of a pilot scale metering roll coater at CB coatweights (when dry) in the range 3.7 to 7.4 g m $^{-2}$. The CF paper utilised acid-washed dioctahedral monimorillonite clay as the capsule rupture. This CB coating composition was applied to the uncoated surface of commercially a gelatinized starch binder and ground cellulose fibre floc as an agent for preventing premature micro active colour developing ingredient. The finished microcapsule dispersion was formulated into a conventional CB coating composition using

The resulting paper was subjected to the following tests:

Calender Intensity (CI) Test

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conventional acid-washed montmorilionite colour developer coated paper, passing the superimposed strips through a laboratory calender to rupture the capsules and thereby produce a colour on the colour This involved superimposing a strip of the microcapsule-coated paper under test onto a strip of

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doveloper strip, measuring the reflectance of the thus-coloured strip (t) and expressing the result $\ell(\omega)$ as a porcentage of the reflectance of an unused control colour developer strip (t₀). Thus the lower the calender intensity value $\ell(\omega)$, the more intense the developed colour.

calendering, the sample being kept in the dark in the interim. Measurements were made both after two minutes and after forty-eight hours, so as to allow for the effect of additional colour development with time. In each case the calender intensity value is indicative of the ability of the microcapsule-coated paper to The reflectance measurements were cione both two minutes after calendering and forty-eight hours after

give rise to a good copy image.

Extended Ram Tost

stack of twenty CFB sheets of each sample was placed under a hydraulic ram and subjected to a nominal ram pressure of 1724 kPa (250 p.s.l) for 30 minutes. The extent of discolouration was assessed visually This is intended to simulate the effect of post-printing discolouration (as described earlier). A

ii. Visual Examination After Printing
This needs no further explanation.

Discolouration on Storage Tests

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Contact Storage
 A stack of twenty CFB sheets of each sample, all with their CF surfaces uppermost, were placed under a 2 kg weight in an oven at 40°C for 3 weeks. A second stack was similarly tested at 60°C for 3 weeks. The extent of discolouration on the CF surfaces was assessed visually.

Singlo CFB should of each sample were placed in ovens under the following conditions, which are believed to simulate the effect of extended storage prior to use in various parts of the world, particularly those with hot climates where discolouration on storage is most problematical.

45 minutes at 150°C

3 days at 32°C and 90% relative humidity

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3 weeks =

3 weeks at 40°C

3 weeks at 60°C

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Again, the extent of discolouration on the CF surfaces was assessed visually. The results of catender intensity tests are set out in Table 1 below:

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Table 1

66.9	io d
	68.2
	68.4
	70.2
	66.0
	67.1
	68.4
	69.6
	72.4
	67.3
	69.0
	67.9
	70.6
	72.7
	2 min.
	Calender Intensity

RSO composition, despite having a lower concentration of dissolved chromogen. This indicates that the that in general the RSO; EHC compositions give similar calender intensity results to those of the 100% Exact comparisons are difficult because of the different dry CB coatweights obtained, but it will be seen

indeed improves copy intensity. inclusion of a proportion of EHC does not have any unacceptable effects on copy-forming capability, and

one week and four weeks after printing. The fact that the extended ram tests were consistent with those for than for the 3:1 RSO:EHC composition. This result was confirmed by examination of 5500 m reels of each either of the compositions containing EHC. The discolouration was lower for the 1:1 RSO:EHC: composition discolouration behaviour. CFB test paper which had been printed on a Muller-Martini four-colour press, examination being carried out paper which had actually been printed shows that the extended ram test is a good predictor of post-print The extended ram test indicated a higher level of discolouration for the 100% RSO composition than for

all conditions for the compositions containing EHC than for the 100% RSO composition. The discolouration was lower for the 1:1 RSO:EHC composition than for the 3:1 RSO:EHC composition. In the contact storage and accelerated ageing tests, the extent of sheet discolouration was lower under

Example 2

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control having exactly the same total colour former concentration (5.0%) as the solvent composition according to the invention. The procedure was as described in Example 1, except that in the final coating This again illustrates the use of a 1:1 RSO:EHC solvent composition, but this time with a 100% RSO

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fibre floc. composition, the binder was a mixture of galatinized starch and carboxymethyl cellulose, and the agent for preventing promature microcepsule rupture was a mixture of wheatstarch particles and ground cellulose

The milling times and the results of primary droplet size testing were as set out in table 2a below:

Solvent Composition	Median Droplet Size (µm)	Milling Time (min)	I.Q.D.	% Oversize*
RSO/EHC	3.05	43	2.18	3.1
100% ASO	3.11	53	2.22	3.7

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As defined in Example 1

ã It will be seen that the inclusion of a proportion of EHC resulted in a significantly reduced milling time and minor improvements in IOD and % Oversize values The results of calender intensity tests are set out in Table 2b below:

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58.8	69.8	6.9	
60.7	71.5	6.2	
63.1	73.5	5.6	
64.6	74.3	4.9	
8.78	77.2	4.3	100% RSO
56.6	65.6	6.6	
58.8	67.4	6.0	
58.0	67.0	5.5 ·	
8.08	69.1	5.4	
63.7	72.7	5.0	RSO/EHC ·
48 hou	2 min.		
		(g m ⁻²)	
Intensity	Calender Intensity	Dry CB Coatweight	Solvent Composition

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intensity values at comparable coatweights. It will be seen that the inclusion of a proportion of EHC into the RSO resulted in significantly improved 35

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ŝ 4.8 g m⁻² CB coatweight 100% RSO sample. It indicated a higher level of discolouration for the 100% RSO composition than for the RSO/EHC composition, despite the lower coatweight of the former. This was in discoloration was more marked than it had been in the extended ram test. confirmed by visual examination of test paper which had actually been printed - in this case the difference The extended ram test was carried out only on the 5.4 g m⁻² CB coalweight RSO/EHC sample and the

å all conditions for the composition containing EHC than for the 100% RSO composition in the contact storage and accelerated ageing tests, the extent of sheet discolouration was lower under

8 namely a 2:3 RSO:EHC composition (i.e. 40% RSO). The control solvent composition was 100% RSO. The procedure was as described in Example 1, except that different milling equipment was used and that the final coating composition was formulated as described in Example 2. The total chromogenic material concentration was 6.4% in each case, instead of 5.0%. This illustrates the use of a solvent composition containing less than 50% by weight of vegetable oil,

Milling times and the results of primary droplet size testing were as set out in Table 3a below

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Table 3a

Solvent Composition	Median Droplet Size (µm)	Milling Time (min)	1.0.0.	% Oversize*
ASO/EHC	3.15	55	1.70	1.7
100% RSO	3.20	105	2.12	4.7

As defined in Example 1

It will be seen that the inclusion of a proportion of EHC resulted in a dramatic reduction in milling time and a significant improvement in IOD and % Oversize values. The higher milling times recorded in this Example compared with previous examples are thought to be a consequence of the different milling equipment used.

The results of calender intensity tests are set out in Table 3b below:

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Table 3b

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	100% RSO					RSO/EHC			Solvent Composition
5.2 6.0 7.0	3.9	6.6	6.1	5.4	4.2	3.7		Coatweight (g m ⁻²)	Microcapsule
67.6 67.6 66.7 65.8	72.2	63.5	64.3	65.4	68.1	70.9	2 min.		Calender
57.0 56.0 55.0	61.3	52.6	53.5	54.6	57.6	60.8	48 hour		Calender Intensity

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႘ intensity values, at comparable coatweights. It will be seen that the inclusion of a large proportion of EHC into the RSO resulted in slightly improved

5.2 g m⁻² CB coatweight 100% RSO sample. It indicated a slightly higher level of discolouration for the 100% RSO composition than for the RSO/EHC composition. This was confirmed by visual examination of marked than it had been in the extended ram test. test paper which had actually been printed. As with Example 2, the difference in discolouration was more The extended ram test was carried out only on the 5.4 g m⁻² CB coalweight RSO/EHC sample and the

Accelerated ageing tests were carried out under the following conditions:

(a) 45 minutes at 150°C

(c) 3 days at 60 °C (b) 3 days at 40°C

(d) 3 weeks at 40°C

(e) 3 weeks at 60°C

than the 100% RSO samples. It was found that the RSO/EHC samples discoloured less than those of the 100% RSO samples. Contact storage testing was also carried out, and the RSO/EHC samples showed less discolouration

The procedure was similar to that described in Example 1 above except that encapsulation was carried out on a laboratory scale, and a smaller pilot-plant coater was used, namely a Dixon pilot plant coater. The discolouration was evaluated salely by means of the extended ram test smaller scale of this work precluded full print testing, which requires long reels, and so post-printing This illustrates the use of a range of different vegetable oils and of a range of different latty acid esters.

The vegetable oils used were rapeseed oil (RSO), sunflower oil (SFO), soybean oil (SBO) and corn oil

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had caprylic acid and capric acid as the main acid moieties (c. 61% and c. 18% respectively) but also (CO).

The fatty acid esters used were 2-dihythoxyl coccete (EHC), isopropyl myristate (IPM), methyl oleate (MO), plyceryl incaprylate caprate (GTCC) and polypropylene glycol dicaprylate/caprate (PGCC). The compositions of the MO and PGCC were as described in more detail earlier in this specification. The GTCC accordance with the invention. butyric and caproic acids (c. 2% in total). GTCC is a trifunctional ester and its use is therefore not in contained minor proportions of other acid moleties, principally lauric acid (c. 9%), myristic acid (c. 6%) and

õ The specific solvent compositions were chosen to complement those evaluated in Examples 1, 2, and 3,

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100% CO	100\$	100\$	100\$	1:1	1:1	1:1	1:1	1:1	1:1	1:1
CO	SBO (")	SFO (")	RSO (Control)	CO : EHC	SFO: EHC	SBO: EHC	RSO: PGCC	RSO: GTCC	RSO: MO	RSO: IPM

The mixture of dissolved chromogenic materials and their concentration (5.0%) was in each case as described for the RSO/EHC solvent compositions of Example 2. The encapsulation procedure was likewise as described in Example 1, except that it was carried out on a laboratory rather than pilot-plant scale. The the binder was a mixture of gelatinized starch and carboxymethylcellulose, and the agent for preventing premature microcapsule rupture was a mixture of wheatstarch particles and ground cellulose fibre floc. microcapsules were formulated and coated on to CF paper largely as described in Example 1 except that

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The evaluation testing was generally as described in Example 1, except that no printing was carried out

The results of primary droplet size testing were as set out in Table 4a below:

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	Solvent Composition	Median Droplet Size (μm)	Milling Time (min)	I.Q.D.	% Oversize*
\$	RSO/IPM	3.10	41	1.71	8.0
	RSO/MO	3.04	36	23	0.8
	RSO/GTCC	3.08	32	1.90	1.7
	RSO/PGCC	3.05	31	1.69	0.3
	SB0/EHC	3.18	ಪಿ	1.63	1.0
8	SFO/EHC	3.18	55	1.61	0.6
	CO/EHC	3.18	4 6	1.64	0.7
	100% RSO	3.13	45	1.48	2.0
	100% SFO	3.12	8	1.82	1.8
	100% SBO	3.14	\$	- 96	2.6
8	100% CO	3.15	50	1.88	2.1

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It will be seen that in each case, the introduction of fatty acid ester gave improved results in some or all tests compared with the corresponding pure vegetable oil. Whilst the 100% RSO had an exceptionally low IQD, it gave worse % Oversize results and longer milling times than when mixed with fatty acid ester.

comparable to the highest of the IQD values for the pure vegetable oils. Its % oversize value was higher than for the mono- and di-ester blends. The mixture of RSO and GTCC required a relatively short milling time, but its IQD value was

equipment on the same base paper, they are assumed to be similar. not measured, but since all were to the same target value, and were applied using the same coating The results of calender intensity testing are set out in Table 4b below. Microcapsule coatweights were

Table 4b

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100% CO	100% SBO	100% SFO	100% RO	COÆHC	SFO/EHC	SBO/EHC	RSO/PGCC	RSO/GTCC	RSO/MO	RSO/IPM		Solvent Composition
75.3	76.2	79.4	74.7	69.3	73.0	71.6	77.3	78.9	70.1	72.8	2 min.	Calender
65.8	68.2	71.2	65.1	60.3	64.5	62.3	68.3	67.2	64.2	63.1	8 hours	Calender Intensity

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မွ gave a more intense colour than the 100% vegetable oil compositions, but that RSO/GTCC and RSO/PGCC were less intense. After 48 hours development, the pattern was similar, although the RSO/GTCC and RSO/PGCC compositions were now of comparable intensity to the 100% vegetable oil composition. It is vegetable oils as discussed earlier. factor in the RSO/GTCC results, in addition to the chemical similarity of glyceryl esters and natural thought that the relatively poor performance of the RSO/PGCC composition may have been due to the presence of small quantities of desensitizing impurities as discussed earlier: This may also have been a It will be seen that after 2 minutes development, most of the compositions according to the invention

samples before and after compression with the ram. The wave length of light used was 600 nm. The results were as set out in Table 4c below: In the extended ram test, an Eirepho reflectance tester was used to measure the reflectance of the

Table 4c

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100% CO	100% SBO	100% SFO	100% RO	CO/EHC	SFO/EHC	SBO/EHC	RSO/PGCC	RSO/GTCC	RSO/MO	RSO/PM		Solvent Composition
89.8	89.9	90.7	90.0	91.0	90.9	91.2	91.0	90.7	90.9	91.1	Before	Reflecta
91.8	92.4	92.3	92.0	92.6	92.3	92.6	92.6	92.4	92.3	92.4	After	Reflectance (%)
2.0	2.5	1.6	2.0	1.6	1.4	1.4	1.6	1.7	1.4	<u>ا</u> ن		Difference

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As dofined in Example 1

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It will be soon that all the 100% vegotiable oil samples showed greater discolouration in the extended ram tost then the corresponding vegotable cilifatty acid ester compositions, although in the case of sunflower oil, the difference was not large. The values for RSO/PGCC and RSO/GTCC were intermediate between the pure oil and the cilimone-functional ester values.

1:1

GNO: EHER

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CNO: EHC CSO:MIS

vegetable oil/fatty.acid ester samples, with the exception of the RSO/GTCC sample, which was better than 100% RSO but comparable to the other 100% vegetable oils. In the contact storage test, the 100% vogetable oil samples showed worse discolouration than the

In the accelerated ageing test, no significant discolouration was observed for any of the samples after 4 wooks at 32 °C and 80% RH.

Example 5

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This illustrates the use of a solvent composition containing a smaller proportion of vegetable oil than in previous examples, namely a 1:3 blend of RSO and EHC (i.e. 25% RSO). The procedure was as described in Example 2, although no 100% RSO control was run.

8 Coulter Counter) was 40 minutes, the percentage of "oversize" droplets, as defined previously, was 2.5% and the IQD value was 1.69. All of these values are comparable with values obtained in previous examples which domonstrates that a 1:3 bland of RSO and EHC gives comparable benefits to those obtained with The milling time required to achieve the target median droplet size of 3.2 + 0.2 µm (as measured by a

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GTEH is glyceryl tris (2-ethylhexanoate).

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performance in a vegetable oil/fatty acid ester solvent functional ester was included in order to evaluate its use is not within the invention as defined, this tri5

100% 100% CSO 100\$ GNO

CNO

Notes

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1:1 1:1 1:1 1:1

RSO: EHC RSO: GTEH

7 (control)

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(see note 1)

100% RSO

The results of calender intensity tests are set out in Table 5 below:

Solvent	Dry CB	Calender	Calender Intensity
Composition	(g m ³)	2 min.	48 hour
RSO/EHC	4.0	73.2	64.8
	- 5.0	70.0	61.3
1:3	5.8	69.5	60.4
	6.6	68.0	59.0
_	6.8	65.5	55.3

compositions according to the invention. Those values are likewise comparable to those obtained with papers utilizing earlier-exemplified

utilizing earliar-exemplified compositions according to the invention. Visual exemination of the paper after printing also demonstrated the comparability of the 1.3 RSO/EHC paper and other papers according to the The extended ram test also gave a degree of discolouration comparable to that shown with papers

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Example 6

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This illustrates the use of a further three vegetable cits, namely groundnut cit (GNO), coconut cit (CNO) and cottonsped cit (CSO), and a further two esters (EHEH and MIS). The procedure was generally as described in Example 1 except that (a) it was carried out on a laboratory scale (b) the chromogenic material bland was a 5% total concentration mixture of CVL, a green fluoran, a black fluoran and a red bis-indolyt phthalide, and (c) the agent for preventing premature microcapsule rupture was a mixture of wheatstarch particles and ground cellulose fibre floc.

The specific solvent compositions evaluated were as follows:

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Solvent	Dry CB	Calender	Calender Intensity
composition	(d m ₋₁)	2 min.	48 hour
RSO/EHC	4.0	73.2	64.8
	5.0	70.0	61.3
1:3	5.8	69.5	60.4
	6.6	68.0	59.0
	6.8	65.5	55.3

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time data was obtained on this occasion because of problems with the milling equipment used.

The results of primary droptet size testing were as set out in Table 6a below. No meaningful milling

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but was included in this evaluation to assist assessment This composition was exemplified in previous Examples,

of the performance of the oils and esters being evaluated

for the first time.

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	% Oversize
GNO/EHEH 3.2 1.6	0.6
3.2	. ن
	0.5
3.2	2.2
3.2	1.5
	1.6
3.2	1.7
	2.0
100% CNO 3.2 1.8	2.6

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ş It will be seen that the oil/ester mixtures gave rise to lower LQ.D. values and % oversize values than the oils alone, with the exception of the RSO/GTEH bland, which is of course not according to the invention.

Table 6b below: The results of calender intensity testing (the mean of three determinations in each case) are set out in

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Table 6b

67.1	71.9	4.7	100% CNO
63.8	68.8	4.4	100% CSO
68.6	73.8	<u>ئ</u>	100% GNO
62.8	67.6	4.5	100% RSO
59.8	62.3	4.7	RSO/EHC
64.3	69.1	4.3	RSO/GTEH
58.3	63.1	5.3	CNO/EHC
60.1	64.3	4.7	CSO/MIS
60.6	64.3	4.2	GNO/EHEH
48 hour	2 min		
Intensity	Calender Intensity	Dry CB Coatweight (g m ⁻²)	Solvent Composition

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It will be seen that the cildester mixture samples gave rise to a more intense colour than the cils alone, with the execution as before, of the RSO/GTEH bland.

with the exception, as before, of the RSO/GTEH blend.

o in the extended ram test, an Eriopho relicctance testor was used to measure the relicctance of the samples before and story compression with the ram. The wave length of light used was 800 nm. The results were as set out in Table 6c below:

Table 6c

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Solvent Composition Reflectance (%) Difference											
ctance (%) After 91.5 90.8 90.8 91.0 91.1 90.4 91.1 90.7	100% CNO	100% CSO	100% GNO	100% RSO	RSO/EHC	RSO/GTEH	CNO/EHC	CSO/MIS	GNO/EHEH		Solvent Composition
	91.6	91.6	91.6	91.3	91.8	91.7	91.6	92.0	92.1	Before	Reflecti
0.6 1.2 0.7 0.7 0.7 0.7 0.9 0.5 0.8	91.1	90.7	91.1	90.4	91.1	91.0	90.9	80.8	91.5	Aftor	ance (%)
	0.5	0.9	0.5	0.9	0.7	0.7	0.7	1.2	0.6		Difference

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40 It will be seen that no clear trend emerges. Possibly this is a consequence of the relatively small differences in reflectance observed in this experiment compared with those observed in Example 4. After accolorated agoing testing for 1 week at 32°C and 90% relative humidity, the GNO/EHEH sample

showed the less discolouration, followed by the RSO/EHC sample, 100% RSO and 100% GNO. The remaining samples all suffered from discolouration to about the same extent. In a separate sat of tests for 3 weeks at 40°C, all the samples showed little discolouration. On testing for 3 weeks at 60°C, all the vegetable oilestor mixture samples showed less discolouration than the 100% vegetable oil samples, which was the best of the samples on test. In the contect storage test, 100% CNO again performed best, followed by the vegetable oil/ester

In the contact storage test, 100% CNO again performed best, lotiowed by the vegetable overster mixture samples and then the remaining 100% vegetable oil samples. The RSO/GTEH sample was the worst of the vegetable oil/ster mixture samples.

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It is thought that the unexpectedly good performance of the 100% coconut oil sample compared with other 100% oil samples is a consequence of the fact that coconut oil solidifies at around ambient temperature, and therefore perhaps flows less freely and hence produces less undesired colouration.

Example 7

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This illustrates the use of triphenylmethane carbinol or carbinol derivative chromogenic materials in the present solvent composition.

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The solvent composition in each case was 1:1 RSO:EHC, with a 100% RSO control. The chromogenic materials were:

$$\begin{array}{c} CH^{0} \\ CH^{0} \\ CH^{0} \\ \end{array}$$

(Example 1 of European Patent Application No. 234394A) and

where X is a mixture of -OH and -OCH₃

(Example 2 of European Patent Application No. 303942A).

A small proportion (less than 2%) of a dialky/naphthalene was present as an impurity in the case where chromogenic material (1) was used.

The milling times and the results of primary droplet size testing were as set out in Table 7 below:

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Table 7

Solvent Composition (Chromogen No.)	Median Droplet Sizo (ит)	Milling Time (min) 1.Q.D.	I.Q.D.	% Oversize*
RSO/EHC (1)	3.19	ಪಿ	1.81	3.0
100% RSO (1)	3.17	. 51	2.35	6.1
RSO/EHC (2)	3.15	ŝ	1.58	0.7
100% RSO (2)	3.13	38	1.98	3.7
. As defined in Example 1	91			

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It will be seen that the solvent compositions according to the invention both gave significantly better LQ.D. and % oversize results than the respective controls. The milling time data is contradictory.

Example 8

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This illustrates the use of the present solvent composition with an encapsulation system relying on in situ polymerisation of aminoplast precondensate for microcapsule wall formation rather than on coacervation of gelatin and other colloids (as in the case of the previous Examples). The aminoplast encapsulation system used is disclosed in full in U.S. Patent No. 4105823.

as a control, using a 100% RSO solvent composition. The solvent composition was a 50:50 mixture of RSO and EHC. A parallel experiment was carried out

chromogenic material solution was similar to that used in Example 6). The resulting emulsion was then acrylic acid contont of 42% by weight ("R144" supplied by Allied Colloids Limited, of Bradford, England) were mixed with 1011 g water, and the mixture was held at 50°C by means of a water bath, 65 g of 20% Warloy, England) were added. The resulting mixture was held in the water bath for 40 minutes before being removed. 243 g of water was added and 1232 ml of chromogenic material solution were added (the milled as described in previous Examples, except that the target droptet size was around 5 um. solids content urea-formaldehyde precondensate ("BC777" supplied by British Industrial Plastics Limited of 274 g of a 20% solids content aqueous dispersion of an acrylic acid/acrylamide copolymer having an

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The milling times and the results of primary droplet size testing were as set out in Table 8 below:

Table 8

* Defined as droplets of diameter greater than 8µm (this different standard, compared with previous Examples, is a consequence of the different encapsulation system being used).

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values than the control. It will be seen that the solvent composition according to the invention gave better I.Q.D. and oversize

Claims

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- A solvent composition for use in pressure-sensitive copying paper and comprising vegetable oil, characterized in that the solvent composition also comprises a proportion of a mone-or diffunctional ester of a non-aromatic mone-carboxylic acid having a saturated or unsaturated straight or branched hydrocarbon chain with at least throe carbon atoms in the chain.
- A solvent composition as claimed in claim 1 whorein the ester is a fatty acid ester or a synthesized fatty acid oster.

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3. A solvent composition as claimed in claim 2 wherein the ester is 2-ethylhexyl cocoate or isopropyl

4. A solvent composition as claimed in claim 1 wheroin the ester is a naturally-occurring lipid or a synthesized such tipid.

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- 5. A solvent composition as claimed in claim 4 wherein the ester is 2-ethylhexyl-2-ethylhexanoate
- A solvent composition as claimed in any preceding claim wherein the vegetable oil is rapeseed oil, soya bean oil, sunflower oil, or corn oil.
- A solvent composition as claimed in any preceding claim wherein the composition is composed substantially entirely of vegetable oil(s) and mono- or di-functional ester(s) as defined in any preceding

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- A solvent composition as claimed in claim 7 wherein the weight ratio of vegetable oil:ester is in the range 1:3 to 3:1.
- A solvent composition as claimed in claim 7 or 8 wherein the vegetable oil is rapeseed oil and the ester
- 10. Pressure-sensitive copying paper comprising a solvent composition as claimed in any preceding claim

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EUROPEAN SEARCH REPORT

	DOCUMENTS CONSIDERED TO BE RELEVANT		
Category	Citation of document with indication, where appropriate, of relevant passages	Relayant to chila	CLASSIFICATION OF THE APPLICATION (Int. CLS.)
×	EP-A-0 390 432 (JUJO PAPER CO., LTD)	1,2,6,10	84195/165
	* page 2, line 35 - line 44 * * page 2, line 53 - line 55 *		
, , o	EP-A-0 024 898 (MONSAMID EUROPE S.A.) * abstract * * page 7, line 18 - page 8, line 6 * * page 8, line 22 - page 9, line 7 *	1-6, 10	
>	ED-A-0 247 816 (FIUI PHOTO FILM CO., LTD.) " claims 1,2" " page 5, line 63 - page 6, line 7"		
>	DE-A-2 618 264 (NCR CORP.)	_	
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CATEGORY OF CITAD DOCUMENTS

X perfoculty relevant if than the

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D: forcement their is the application because of their stars pasted healthy, corresponding forces and

The present exacts report has been drawn up for all claims

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O2 SEPTEMBER 1992

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Application Number